The structure for the search was:

The benzophenone gave 139 hits. These did not seem relevant so I did a search for the structure and (THYROID OR THRYOMIMETIC OR ?THYRONINE). Four hits came up and they are at the bottom of this search.

ANSWER 1 OF 21 HCAPLUS COPYRIGHT 1999 ACS L9

AN 1999:9803 HCAPLUS

ΤI Preparation of phenoxyakanoates as thyroid hormone receptor .beta. agonists

IN Scanlan, Thomas S.; Chellini, Grazia; Yoshihara, Hikari; Apriletti, James;

Baxter, John D.; Ribeiro, Ralff C. J.

PA The Regents of the University of California, USA

PCT Int. Appl., 45 pp. so

CODEN: PIXXD2

DT Patent

LA English

AB R30Z1CR1R2Z20(CH2)nCO2R [I; R = H or (cyclo)alkyl; R1,R2 = H or alkyl; 1 of R1, R2 = H and the other = OH; R1R2 = O; R3 = H, (cyclo)alkyl, acyl; Z1 = (un)substituted 1,4-phenylene; Z2 = (un)substituted 3,5-dimethyl-4,1phenylene] were prepd. Thus, 4-bromo-2-isopropylanisole was condensed with 2,6-dimethyl-4-methoxybenzaldehyde (prepn. each given) and the product converted in 4 steps to title compd. II. Data for biol. activity of I were given.

IT 218431-15-3P

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of phenoxyakanoates as thyroid hormone receptor .beta. agonists)

RN 218431-15-3 HCAPLUS

CN INDEX NAME NOT YET ASSIGNED

IT 211110-65-5P 218431-12-0P 218431-13-1P

218431-14-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
 (prepn. of phenoxyakanoates as thyroid hormone receptor .beta.
 agonists)

RN 211110-65-5 HCAPLUS

CN Benzenemethanol, 4-methoxy-.alpha.-[4-methoxy-3-(1-methylethyl)phenyl]-

2,6-

dimethyl- (9CI) (CA INDEX NAME)

RN 218431-12-0 HCAPLUS

CN INDEX NAME NOT YET ASSIGNED

RN 218431-13-1 HCAPLUS

CN INDEX NAME NOT YET ASSIGNED

RN 218431-14-2 HCAPLUS

CN INDEX NAME NOT YET ASSIGNED

L9 ANSWER 2 OF 21 HCAPLUS COPYRIGHT 1999 ACS

AN 1998:617873 HCAPLUS

DN 129:302827

TI An efficient substitution reaction for the preparation of thyroid hormone analoges

AU Yoshihara, Hikari A. I.; Chiellini, Grazia; Mitchison, Timothy J.; Scanlan, Thomas S.

CS Department of Cellular and Molecular Pharmacology, University of California, San Francisco, CA, 94143-0450, USA

SO Bioorg. Med. Chem. (1998), 6(8), 1179-1183 CODEN: BMECEP; ISSN: 0968-0896

PB Elsevier Science Ltd.

DT Journal

LA English

AB The substitution of the sterically hindered carbon of the potent thyroid hormone agonist, GC-1, was effected by a reaction based on the solvolysis of the benzylic hydroxyl group. The reaction was found to proceed in

high

yield with a variety of nucleophiles including alcs., thiols, allyl silanes and electron-rich arom. compds., providing a convenient route to the synthesis of new thyroid hormone analogs.

IT 211110-65-5

RL: RCT (Reactant)

(prepn. of thyroid hormone analoges via substitution reaction)

RN 211110-65-5 HCAPLUS

CN Benzenemethanol, 4-methoxy-.alpha.-[4-methoxy-3-(1-methylethyl)phenyl]-

2,6-

dimethyl- (9CI) (CA INDEX NAME)

IT 214544-37-3P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of thyroid hormone analoges via substitution reaction)

RN 214544-37-3 HCAPLUS

CN Benzene, 2-[ethoxy[4-methoxy-3-(1-methylethyl)phenyl]methyl]-5-methoxy-

1,3-

dimethyl- (9CI) (CA INDEX NAME)

L9 ANSWER 3 OF 21 HCAPLUS COPYRIGHT 1999 ACS

AN 1998:435316 HCAPLUS

DN 129:157050

TI A high-affinity subtype-selective agonist ligand for the thyroid hormone receptor $\frac{1}{2^{N}}\frac{1}{N}$

AU Chiellini, Grazia; Apriletti, James W.; Yoshihara, Hikari Al; Baxter, John

D.; Ribeiro, Ralff C. J.; Scanlan, Thomas S.

CS Department of Pharmaceutical Chemistry and Cellular & Molecular Pharmacology, University of California, San Francisco, CA, 94143-0446,

USA

SO Chem. Biol. (1998), 5(6), 299-306 CODEN: CBOLE2; ISSN: 1074-5521

PB Current Biology Ltd.

DT Journal

LA English

AB Thyroid hormones regulate many different physiol. processes in different tissues in vertebrates. Most of the actions of thyroid hormones are mediated by the thyroid hormone receptor (TR), which is a member of the nuclear receptor superfamily of ligand-activated transcription regulators.

There are two different genes that encode two different TRs, TR.alpha.

and

TR.beta., and these two TRs are often co-expressed at different levels in different tissues. Most thyroid hormones do not discriminate between the two TRs and bind both with similar affinities. The authors have designed and synthesized a thyroid hormone analog that has high affinity for the

TRs and is selective in both binding and activation functions for TR.beta.

over TR.alpha.. The compd., GC-1, was initially designed to solve synthetic problems that limit thyroid hormone analog prepn., and contains several structural changes with respect to the natural hormone 3,5,3'-triiodo-L-thyronine (T3). These changes include replacement of

three iodines with Me and iso-Pr groups, replacement of the biaryl ether linkage with a methylene linkage, and replacement of the amino-acid sidechain with an oxyacetic-acid sidechain. The result of this study

that GC-1 is a member of a new class of thyromimetic compds. that are more $% \left(1\right) =\left(1\right) +\left(1\right)$

synthetically accessible than traditional thyromimetics and have potentially useful receptor binding and activation properties. The TR.beta. selectivity of GC-1 is particularly interesting and suggests that

GC-1 might be a useful in vivo probe for studying the physiol. roles of the different thyroid hormone receptor isoforms.

211110-65-5P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (design and synthesis of high-affinity subtype-selective agonist liquid

for thyroid hormone receptor)

RN 211110-65-5 HCAPLUS

the

show

CN Benzenemethanol, 4-methoxy-.alpha.-[4-methoxy-3-(1-methylethyl)phenyl]-2,6-

dimethyl- (9CI) (CA INDEX NAME)

- L9 ANSWER 4 OF 21 HCAPLUS COPYRIGHT 1999 ACS
- AN 1998:432999 HCAPLUS
- DN 129:245014
- TI Synthesis and biological activity of 2,3-benzopyrone analogs
- AU Ji, Xiaoshen; Liang, Xiaotian
- CS Department of Clinical Pharmacy, General Hospital of Air Force, PLA, Beijing, 100036, Peop. Rep. China
- SO Yaoxue Xuebao (1998), 33(1), 72-74 CODEN: YHHPAL; ISSN: 0513-4870
- PB Chinese Academy of Medical Sciences, Institute of Materia Media
- DT Journal
- LA Chinese
- AB The Friedel-Crafts reaction was taken place with some replacement Ph acetic acid or its Me ester and vanillin reactants in the condition of Ac2O/ZnCl2. Two compds. showed obvious activities on the potassium channel and anticancer screen.
- IT 213138-34-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (synthesis and biol. activity of 2,3-benzopyrone analogs)

- RN 213138-34-2 HCAPLUS
- CN Benzeneacetic acid, 4-(acetyloxy)-2-[(acetyloxy)[4-(acetyloxy)-3-methoxyphenyl]methyl]-3,5-dimethoxy-, methyl ester (9CI) (CA INDEX NAME)

- L9 ANSWER 5 OF 21 HCAPLUS COPYRIGHT 1999 ACS
- AN 1997:667252 HCAPLUS
- DN 127:293323
- TI Synthesis and Chemistry of CF3C6F4OC6F4 Group 14/16 Derivatives
- AU Krumm, Burkhard; Kirchmeier, Robert L.; Shreeve, Jean'ne M.
- CS Department of Chemistry, University of Idaho, Moscow, ID, 83844-2343, USA
- SO Inorg. Chem. (1997), 36(23), 5222-5230 CODEN: INOCAJ; ISSN: 0020-1669
- PB American Chemical Society
- DT Journal
- LA English

is

- OS CASREACT 127:293323; CJACS
- AB Reactions of 4'-CF3C6F4OC6F4Li, generated in situ, with elements of group 16 (S, Se, Te) lead to CF3C6F4OC6F4SH (2), (CF3C6F4OC6F4Se)2 (3), and (CF3C6F4OC6F3Te)2 (4)/(CF3C6F4OC6F3)2Te (4a). The phenol deriv. CF3C6F4OC6F4OH (1) is obtained by reaction of CF3C6F4OC6F4Li with B(OMe)3/H2O2. The reaction of CF3C6F4OC6F4Li with trimethylsilyl chloride
 - or trimethyltin chloride gives CF3C6F4OC6F4XMe3 (X = Si (5), Sn (6)). Oxidn. of 2 in the presence of bromine results in the formation of (CF3C6F4OC6F4S)2 (7) and CF3C6F4OC6F4SO2Br (8). Mixed perfluoroaryloxo/thio ethers CF3C6F4OC6F4SC6F4R (R = NO2 (9), CN (10),
- CF3 (11)) and CF3C6F4OC6F4SC5F4N (12) are obtained upon reaction of 2 with excess C6F5R and pentafluoropyridine in the presence of K2CO3. With 4-C6F5OC6F4NO2, a mixt. of (2-CF3C6F4OC6F4S)(4-C6F5O)C6F3NO2 (13) and 9
- formed. Reaction of excess 2 with C6F5R gives the 2,4,6-substituted benzenes (CF3C6F4OC6F4S)3C6F2R (R = NO2 (14), CN (15)). The trimethylsilyl ether CF3C6F4OC6F4OSiMe3 (16) is prepd. from the reaction of 1 with hexamethyldisilazane. 16 Is a convenient reagent for the prepn.
- of the aryl ethers CF3C6F40C6F40C6F4R (R = NO2 (17), CN (18)) and CF3C6F40C6F40C5F4N (19) upon reaction with C6F5R and C5F5N. The secondary
 - alcs. CF3C6F4OC6F4CH(C6H5)OH (20) and CF3C6F4OC6F4CH(C6F5)OH (21) are synthesized by the reactions of 5 with benzaldehyde and

pentafluorobenzaldehyde in the presence of tetrabutylammonium fluoride as a catalyst. In the synthesis of 21 the byproduct

CF3C6F4OC6F4CH(C6F5)OC6F4CHO is also formed and isolated.

IT 197150-25-7P 197150-26-8P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (prepn. of)

RN 197150-25-7 HCAPLUS

CN Benzenemethanol, 2,3,4,5,6-pentafluoro-.alpha.-[2,3,5,6-tetrafluoro-4-[2,3,5,6-tetrafluoro-4-(trifluoromethyl)phenoxy]phenyl]- (9CI) (CA INDEX NAME)

RN 197150-26-8 HCAPLUS

CN Benzaldehyde, 2,3,5,6-tetrafluoro-4-[(pentafluorophenyl)]2,3,5,6-tetrafluoro-4-

(trifluoromethyl)phenoxy]phenyl]metho

xy] - (9CI) (CA INDEX NAME)

L9 ANSWER 6 OF 21 HCAPLUS COPYRIGHT 1999 ACS

AN 1997:271246 HCAPLUS

DN 126:317282

TI Synthesis and hypolipidemic activity of diesters of arylnaphthalene lignan

and their heteroaromatic analogs

AU Kuroda, Tooru; Kondo, Kazuhiko; Iwasaki, Tameo; Ohtani, Akio; Takashima, Kohki

CS Res. Lab. Tanabe Seiyaku Co., Ltd., Osaka, 532, Japan

SO Chem. Pharm. Bull. (1997), 45(4), 678-684

CODEN: CPBTAL; ISSN: 0009-2363

PB Pharmaceutical Society of Japan

DT Journal LA English GI

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB A series of arylnaphthalene lignan diesters (I) (R1 = Me, Et, CHMe2, C6H13, C10H21, CH2Ph, CH2CH2OMe, CH2CH2NEt2.HCl, CH2CH2-4-morpholine.HCl, 3-pyridyl.HCl, cyclohexylmethyl, CH2Ph; R2 = Me, Et, CHEt2, C6H13, cyclohexylmethyl, CH2Ph)) and their heteroarom. analogs II (R3 = Me, Et) and III (R4 = SO2Ph, H) were synthesized and evaluated for hypolipidemic activity. The diesters with modifications at C-3 showed excellent hypocholesterolemic and high-d. lipoprotein (HDL) cholesterol-elevating activities. Structure-activity anal. indicated that I (R1 = 2-pyridylmethyl.HCl, R2 = Me) has the optimum activity.

IT 104756-71-0

RL: RCT (Reactant)

(synthesis and hypolipidemic activity of diesters of arylnaphthalene lignan and their heteroarom. analogs)

RN 104756-71-0 HCAPLUS

CN Benzenemethanol, 6-(dimethoxymethyl)-.alpha.-(3,4-dimethoxyphenyl)-2,3,4-trimethoxy- (9CI) (CA INDEX NAME)

L9 ANSWER 7 OF 21 HCAPLUS COPYRIGHT 1999 ACS

AN 1996:733900 HCAPLUS

DN 126:31215

TI Efficient Synthesis of 1-Aryl-3,4-dihydro-4-hydroxynaphthalene: Application to the Stereocontrolled Synthesis of (.+-.)-Isopicropodophyllin and (.+-.)-Isopodophyllotoxin

AU Kuroda, Tooru; Takahashi, Masami; Kondo, Kazuhiko; Iwasaki, Tameo

CS Pharmaceutical Development Research Laboratory, Tanabe Seiyaku Co. Ltd., Osaka, 532, Japan

SO J. Org. Chem. (1996), 61(26), 9560-9563 CODEN: JOCEAH; ISSN: 0022-3263

PB American Chemical Society

DT Journal

LA English

OS CJACS

GΙ

AB An efficient method for synthesizing naphthalenes I (R1=R2=R3 = OMe, R4 = H; R1,R2 = OCH2O, R3 = H, R4 = OMe) via the acid-catalyzed reaction of acetoxyaldehydes with di-Me maleate is presented. Also, the authors have shown that I (R1,R2 = OCH2O, R3 = H, R4 = OMe) can be transformed to (.+-.)-isopicropodophyllin and (.+-.)-isopodophyllotoxin via stereocontrolled hydrogenations.

IT 131924-17-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (synthesis of (.+-.)-isopicropodophyllin and (.+-.)-isopodophyllotoxin via stereocontrolled hydrogenation of aryldihydrohydroxynaphthalenes)

RN 131924-17-9 HCAPLUS

CN Benzaldehyde, 2-[(acetyloxy)(3,4-dimethoxyphenyl)methyl]-3,4,5-trimethoxy-

(9CI) (CA INDEX NAME)

L9 ANSWER 8 OF 21 HCAPLUS COPYRIGHT 1999 ACS

Ι

- AN 1995:959433 HCAPLUS
- DN 124:105580
- TI Arylnaphthalene lignans as novel series of hypolipidemic agents raising high-density lipoprotein level
- AU Iwasaki, Tameo; Kondo, Kazuhiko; Nishitani, Takashi; Kuroda, Tooru; Hirakoso, Kazuyuki; Ohtani, Akio; Takashima, Kohki
- CS Res. Lab. Tanabe Seiyaku Co., Ltd., Osaka, 532, Japan
- SO Chem. Pharm. Bull. (1995), 43(10), 1701-5 CODEN: CPBTAL; ISSN: 0009-2363
- DT Journal
- LA English
- AB A series of arylnaphthalene lignans were prepd. and tested for hypolipidemic activity. The most potent compd. (TA-7552) not only reduced

serum cholesterol, but also increased high-d. lipoproteins cholesterol in rats. The ED of TA-7552 is 100-fold less than that of cholestyramine. Structure-activity relations are discussed.

104756-71-0P IT

> RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (in prepn. of arylnaphthalene lignans as hypolipidemic agents increasing high-d. lipoproteins)

RN 104756-71-0 HCAPLUS

CN Benzenemethanol, 6-(dimethoxymethyl)-.alpha.-(3,4-dimethoxyphenyl)-2,3,4trimethoxy- (9CI) (CA INDEX NAME)

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L9
    ANSWER 9 OF 21 HCAPLUS COPYRIGHT 1999 ACS
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AN 1995:794873 HCAPLUS

DN 123:198645

ΤI Preparation of balanoids as protein kinase C inhibitors

IN Hall, Steven Edward; Ballas, Lawrence M.; Kulanthaivel, Palaniappan; Boros, Christie; Jiang, Jack B.; Jagdmann, Gunnar Erik, Jr.; Lai, Yen-Shi;

Biggers, Christopher K.; Hu, Hong; et al.

PA Nichols, Gina M., USA; Sphinx Pharmaceuticals Corporation

SO PCT Int. Appl., 559 pp.

CODEN: PIXXD2

DΤ Patent

English T.A

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		W:	ΑT,	AU,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CZ,	DE,	DK,	ES,	FI,	GB,	HU,
			JP,	KP,	KR,	KZ,	LK,	LU,	LV,	MG,	MN,	MW,	NL,	NO,	NZ,	PL,	PT,	RO,
			RU,	SD,	SE,	SK,	UA,	US,	UZ,	VN								
		RW:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,
			BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	ML,	MR,	NE,	SN,	TD,	TG		
	CA	2157	412		A.	A	1994	0915		C	A 94	-215	7412		1994	0302		
	AU	9462527			A1 19940926			AU 94-62527			19940302							
	ΕP	6872	49		A1 19951220			EP 94-909847			19940302							
		R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙE,	IT,	LI,	LU,	MC,	NL,	PT,
SE																		
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	ZA	9401	478		Α		1995	0905		\mathbf{z}_{i}	A 94	-147	В		1994	0303		
PRAI	US	93-2	5846		199	9303	03											
	WO	94-U	S228	3	199	9403	02											



OS MARPAT 123:198645

GΙ

AB Title compds. [I; A = CH2, NR1, O, S, SO2; B1 = NR2, CH2, O; B2 = CO, CS, SO2; D = NR3 = O, CH2; E = R5, (un)substituted (hetero)arylene; F = CO or CH2; G = R7, cycloalkyl, (un)substituted (hetero)aryl; K = H, alkyl; R = R4, (un)substituted Ph, (hetero)aryl; R1-R4, R7 = H, alkyl, aryl, etc.;

IT 167832-20-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. of balanoids as protein kinase C inhibitors)

RN 167832-20-4 HCAPLUS

CN Benzoic acid, 4-[hydroxy[4-(phenylmethoxy)-3-

[(phenylmethoxy)carbonyl]phen

yl]methyl]-3,5-bis(phenylmethoxy)-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

L9 ANSWER 10 OF 21 HCAPLUS COPYRIGHT 1999 ACS

AN 1991:206825 HCAPLUS

DN 114:206825

TI Preparations of hypolipemic 1-phenyl-2,3-bis(alkoxycarbonyl)-4-hydroxynaphthalenes and their intermediates

IN Iwasaki, Tameo; Nishitani, Takashi; Omizu, Hiroshi; Takahashi, Masami; Oogiku, Ko

Tanabe Seiyaku Co., Ltd., Japan PA so

Jpn. Kokai Tokkyo Koho, 7 pp.

CODEN: JKXXAF

Patent DT

Japanese LA

FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. JP 02300148 PI A2 19901212 JP 89-117955 19890511

MARPAT 114:206825 os

GΙ

Me.

OH
$$CO_2R^1$$
 A CO_2R^2 CO_2R^2

AB A process for the prepn. of the title compds. I (R1, R2 = lower alkyl; R3,

R4 = H, lower alkoxy; R3 and/or R4 = lower alkoxy; ring A may be substituted) or their salts, useful as hypolipemics (no data), by oxidn. of dihydronaphthalenes II or their salts, which may be prepd. by

of 2-(phenylhydroxymethyl)benzaldehydes III (R5 = H, hydroxy-protective group), their di-lower alkyl acetals, or their salts with R10COCH: CHCO2R2,

optionally followed by salt formation, and II or their salts are claimed. 2-(.alpha.-Hydroxy-3,4-dimethoxybenzyl)-3,4,5-trimethoxybenzaldehyde di-Me

acetal (816 mg) in di-Me maleate was added dropwise to CF3CO2H in di-Me maleate at 70.degree. over 2.5 h and the reaction mixt. was further stirred at 70.degree. for 1.5 h to give 330 mg (r-3,t-4)-II (R1 = R2 =

R3 = R4 = OMe, 6, 7, and 8-positions are substituted with OMe). This (600

mg) in dioxane was treated with 2,3-dichloro-5,6-dicyanobenzoquinone under

stirring at 80.degree. for 35 h to give 240 mg I (R1 = R2 = Me, R3 = R4 = OMe, 6, 7, and 8-positions are substituted with OMe).

131924-17-9P 131924-18-0P 133491-26-6P IT 133491-27-7P 133491-28-8P 133491-29-9P

> RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and cyclocondensation of, with dialkyl maleate or fumarate,

phenylhydroxydihydronaphthalenedicarboxylate from)

RN 131924-17-9 HCAPLUS

CN Benzaldehyde, 2-[(acetyloxy)(3,4-dimethoxyphenyl)methyl]-3,4,5-trimethoxy-

(9CI) (CA INDEX NAME)

RN 131924-18-0 HCAPLUS

CN Benzaldehyde, 2-[(3,4-dimethoxyphenyl)methoxymethyl]-3,4,5-trimethoxy-(9CI) (CA INDEX NAME)

RN 133491-26-6 HCAPLUS

CN Benzaldehyde, 2-[(3,4-diethoxyphenyl)hydroxymethyl]-3,4,5-trimethoxy-(9CI) (CA INDEX NAME)

RN 133491-27-7 HCAPLUS

CN Benzaldehyde, 2-[(3,4-dipropoxyphenyl)hydroxymethyl]-3,4,5-trimethoxy-(9CI) (CA INDEX NAME)

RN 133491-28-8 HCAPLUS

CN Benzaldehyde, 2-[(3-ethoxy-4-methoxyphenyl)hydroxymethyl]-3,4,5-trimethoxy-

(9CI) (CA INDEX NAME)

RN 133491-29-9 HCAPLUS

CN Benzaldehyde, 2-[(4-ethoxy-3-methoxyphenyl)hydroxymethyl]-3,4,5-trimethoxy-

(9CI) (CA INDEX NAME)

IT 131924-15-7P 131924-16-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
 (prepn. and deacetalization of)

RN 131924-15-7 HCAPLUS

CN Benzenemethanol, 6-(dimethoxymethyl)-.alpha.-(3,4-dimethoxyphenyl)-2,3,4-trimethoxy-, acetate (9CI) (CA INDEX NAME)

RN 131924-16-8 HCAPLUS

CN Benzene, 1-(dimethoxymethyl)-2-[(3,4-dimethoxyphenyl)methoxymethyl]-3,4,5-

trimethoxy- (9CI) (CA INDEX NAME)

IT 104756-71-0

RL: RCT (Reactant)

(reaction of, in prepn. of hypolipemic dialkyl (alkoxyphenyl)hydroxynaphthalenedicarboxylates)

RN 104756-71-0 HCAPLUS

CN Benzenemethanol, 6-(dimethoxymethyl)-.alpha.-(3,4-dimethoxyphenyl)-2,3,4-trimethoxy- (9CI) (CA INDEX NAME)

L9 ANSWER 11 OF 21 HCAPLUS COPYRIGHT 1999 ACS

AN 1991:81276 HCAPLUS

DN 114:81276

TI Process for preparing 1-hydroxy-4-phenylnaphthalene-2,3-dicarboxylates useful as antihyperlipidemics

IN Iwasaki, Tameo; Ohmizu, Hiroshi; Tsuyoshi, Ohgiku

PA Tanabe Seiyaku Co., Ltd., Japan

SO Eur. Pat. Appl., 17 pp.

CODEN: EPXXDW

DT Patent

LA English

FAN.CNT 1

FAN.CN	N.I.	1									
I	PATENT NO.				KIND DATE			APPLICATION NO.			DATE
-											
PI E	EΡ	37993	35		A:	1 1990	0801	EP	90-100	332	19900116
		R:	AT,	BE,	CH,	DE, DK,	ES, FR	, GB, G	R, IT,	LI, LU	, NL
	CN	10444	156		Α	1990	8080	CN	89-109	562	19891228
2	ZA	90000	77		Α	1990	1031	ZA	90-77		19900105
C	CA	20079	581		A/	A 1990	0727	CA	90-200	7581	19900111
F	HU	53862	2		A2	2 1990	1228	HU	90-173		19900117
I	UA	90489	591		A:	1 1990	0802	AU	90-485	91	19900118
Į	UA	61633	37		B	2 1991	1024				

	JP 0	2275840	A2	19901109	JР	90-15838	19900125
	NO 9	9000381	Α	19900730	NO	90-381	19900126
	SU 1	1831473	A3	19930730	SU	90-4742864	19900126
PRAI	JP 8	39-18587	198901	127			
os	MARE	PAT 114:81276					
GI							

$$R_{\rm R}$$
 CO_2R^1
 CO_2R^2
 R_3
 R_4
 R_5
 R_5
 R_4
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 R_5
 R_6
 R_6
 R_7
 R_7

AB Naphthalene derivs. [I; R = substituent; R1, R2 = alkyl, one of R3 and R4 is H, alkoxy, the other is alkoxy; n = 0-3], useful as hypolipidemic agents (no data), are prepd. by cyclocondensation of benzaldehyde derivs II (R5 = protecting group) with R1O2CC.tplbond.CCO2R2 followed by optional

salt formation. A mixt. of benzaldehyde deriv. III (prepn. given) and MeO2CC.tplbond.CCO2Me in CF3CO2H and C6H6 was heated at 60.degree. to give

77% I [Rn = 6,7,8-(MeO)3, R1 = R2 = Me; R3 = R4 = MeO]. Also prepd. was 22 addnl. I.

IT 104756-71-0

RL: RCT (Reactant)
 (acetylation of)

RN 104756-71-0 HCAPLUS

CN Benzenemethanol, 6-(dimethoxymethyl)-.alpha.-(3,4-dimethoxyphenyl)-2,3,4-trimethoxy- (9CI) (CA INDEX NAME)

IT 131924-17-9P 131924-18-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and cyclocondensation of, with di-Me acetylenedicarboxylate)

RN 131924-17-9 HCAPLUS

CN Benzaldehyde, 2-[(acetyloxy)(3,4-dimethoxyphenyl)methyl]-3,4,5-trimethoxy-

(9CI) (CA INDEX NAME)

RN 131924-18-0 HCAPLUS

CN Benzaldehyde, 2-[(3,4-dimethoxyphenyl)methoxymethyl]-3,4,5-trimethoxy-(9CI) (CA INDEX NAME)

IT 131924-15-7P 131924-16-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
 (prepn. and hydrolysis of)

RN 131924-15-7 HCAPLUS

CN Benzenemethanol, 6-(dimethoxymethyl)-.alpha.-(3,4-dimethoxyphenyl)-2,3,4-trimethoxy-, acetate (9CI) (CA INDEX NAME)

RN 131924-16-8 HCAPLUS

CN Benzene, 1-(dimethoxymethyl)-2-[(3,4-dimethoxyphenyl)methoxymethyl]-3,4,5-

trimethoxy- (9CI) (CA INDEX NAME)

L9 ANSWER 12 OF 21 HCAPLUS COPYRIGHT 1999 ACS

AN 1990:630978 HCAPLUS

DN 113:230978

TI Preparation of 1-(3,4-dialkoxyphenyl)-6,7,8-trialkoxy-4-

hydroxynaphthalene-

2,3-dicarboxylates as hypolipemic agents

IN Suzuki, Takashi; Yamamura, Minehiko; Yamada, Sinichi

PA Tanabe Seiyaku Co., Ltd., Japan

SO Eur. Pat. Appl., 8 pp.

CODEN: EPXXDW

DT Patent

LA English

FAN.CNT 1

PAIN.	-14 T	1							
	PATENT NO.			KIND	DATE		APPLICATIO	N NO.	DATE
ΡI	EР	371484		A 2	19900606		EP 89-1220	10	19891129
	ΕP	371484		A3	19910410				
		R: AT,	BE,	CH, DE	, ES, FR,	GB,	GR, IT, LI,	LU, NL	, SE
	JP	02149546		A2	19900608		JP 88-3033	35	19881129
	CA	2002612		AA	19900629		CA 89-2002	612	19891109
	CN	1043932		Α	19900718		CN 89-1086	52	19891116
	US	5066825		Α	19911119		US 89-4370	65	19891116
	ZA	8908900		Α	19900829		ZA 89-8900)	19891122
	AU	8945513		A1	19900607		AU 89-4551	.3	19891123
	ΑU	613250		B2	19910725				
	DK	8905996		Α	19900530		DK 89-5996		19891128
	NO	8904737		A	19900530		NO 89-4737	,	19891128
	NO	170010		В	19920525				
	NO	170010		С	19920902				
	HU	53060		A2	19900928		HU 89-6312	!	19891129
	HU	204023		В	19911128				

PRAI JP 88-303335 19881129

OS MARPAT 113:230978

GΙ

$$R^{70}$$
 CO_2R^1
 CO_2R^2
 OMe
 OMe

AB The title compds. (I; R1-R7 = alkyl) were prepd. as hypolipemics (no data)

by cyclocondensation of hydroxybenzylbenzaldehyde acetals with acetylenedicarboxylates. Thus, 3,4,5-(MeO)3C6H2CH(OMe)2 (prepn. given) was stirred 30 min at 0.degree. with BuLi in THF after which 3,4-(MeO)2C6H3CHO was added and the whole stirred 2 h at 0-10.degree. to give aldol product II which was refluxed 3 h with MeO2CC.tplbond.CCO2Me

in PhMe contg. 4-MeC6H4SO3H to give I (R1 - R7 = Me).

IT 104756-71-0P 130422-12-7P 130422-13-8P 130422-14-9P 130422-15-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and reaction of, in prepn. of hypolipemic agents)

RN 104756-71-0 HCAPLUS

CN Benzenemethanol, 6-(dimethoxymethyl)-.alpha.-(3,4-dimethoxyphenyl)-2,3,4-trimethoxy- (9CI) (CA INDEX NAME)

RN 130422-12-7 HCAPLUS

CN Benzenemethanol, 6-(dimethoxymethyl)-.alpha.-(4-ethoxy-3-methoxyphenyl)-2,3,4-trimethoxy- (9CI) (CA INDEX NAME)

RN 130422-13-8 HCAPLUS

CN Benzenemethanol, 6-(dimethoxymethyl)-.alpha.-(3-ethoxy-4-methoxyphenyl)-2,3,4-trimethoxy- (9CI) (CA INDEX NAME)

RN 130422-14-9 HCAPLUS

CN Benzenemethanol, .alpha.-(3,4-diethoxyphenyl)-6-(dimethoxymethyl)-2,3,4-trimethoxy- (9CI) (CA INDEX NAME)

RN 130422-15-0 HCAPLUS

CN Benzenemethanol, 6-(dimethoxymethyl)-.alpha.-(3,4-dipropoxyphenyl)-2,3,4-trimethoxy- (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{OPr-n} \\ \text{OH} \\ \text{OMe} \\ \text{OMe} \\ \text{OMe} \\ \end{array}$$

L9 ANSWER 13 OF 21 HCAPLUS COPYRIGHT 1999 ACS

AN 1990:55275 HCAPLUS

DN 112:55275

TI Preparation of phenylnaphthoates and phenylnaphthamides as hypolipemics

PA Tanabe Seiyaku Co., Ltd., Japan

SO Austrian, 17 pp.

CODEN: AUXXAK

DT Patent

LA German

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
ΡI	AT 388372	В	19890612	AT 87-2625	19871008	
	AT 8702625	A	19881115			

OS MARPAT 112:55275

GI

The title compds. [I; A = (un)substituted benzene ring; R1, R2 = C1-4 alkoxy, OR5, NHR5, NR6R7; R3, R4 = H, C1-4 alkoxy; R5 = (un)substituted C1-4 alkyl, C5-10 alkyl, C2-10 alkenyl, C5-8 cycloalkyl, 5- or 6-membered N-heterocyclyl; R6, R7 = H, C1-4 alkyl; R8 = H] and their salts were prepd. as hypolipemics useful for the prevention and treatment of arteriosclerosis, by a cyclocondensation reaction of acetylenedicarboxylates R1COC.tplbond.CCOR2 (II) (R1, R2 as above) with III (R3, R4 as defined) or by esterification or amidation of I (R1 = OH) with R1H. Thus, a mixt. of 1.4 g 1-(3,4-dimethoxyphenyl)-2-methoxycarbonyl-4-benzyloxy-6,7,8-trimethoxy-3-naphthoic acid, 183 mg H2NCH2CHMe2, and 336 mg 1-hydroxybenzotriazole in THF was treated and stirred with 570 mg N,N'-dicyclohexylcarbodiimide for 2 h at 0.degree.

and

12 h at room temp. The intermediate 4-benzyloxy-3-naphthamide was deprotected by stirring 2 h with Pd/C in MeOH, in a H atm. at 3 kg/cm2,

to

give 1.1 g I (R1 = HNCH2CHMe2, R2-R4 = OMe, R8 = H, A = Q). The latter

in

rats reduced total serum cholesterol 60% and increased serum HDL-cholesterol 99%.

IT 104756-71-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and reaction of, in prepn. of hypolipemic)

RN 104756-71-0 HCAPLUS

CN Benzenemethanol, 6-(dimethoxymethyl)-.alpha.-(3,4-dimethoxyphenyl)-2,3,4-trimethoxy- (9CI) (CA INDEX NAME)

L9 ANSWER 14 OF 21 HCAPLUS COPYRIGHT 1999 ACS

AN 1988:630583 HCAPLUS

DN 109:230583

TI Preparation of 4-phenyl-1-naphthol derivatives as hypolipidemic agents

IN Iwasaki, Tameo; Takashima, Koki

PA Tanabe Seiyaku Co., Ltd., Japan

SO Jpn. Kokai Tokkyo Koho, 14 pp.

CODEN: JKXXAF

DT Patent

LA Japanese

FAN. CNT 1

PRAI JP 86-155413 19860701

OS MARPAT 109:230583

GI

AB Title compds. I or II (R1 = H, alkoxycarbonyl; R2 = alkoxycarbonyl; R3, R4

= H, alkoxy, but R3 = R4 .noteq. H; ring A may be substituted) and their salts are prepd. as hypolipidemic agents. A soln. of 204.0 g 2-bromo-3,4,5-trimethoxybenzaldehyde di-Me acetal in THF was treated with BuLi at -70.degree. to -60.degree., then a soln. of 105.5 g

3,4-(MeO)2C6H3CHO in THF was added to give 266 g 2-(3,4-dimethoxy-.alpha.-

hydroxybenzyl)-3,4,5-trimethoxybenzaldehyde di-Me acetal, which was treated with 95 mL MeO2CC.tplbond.CCO2Me and 300 mg p-MeC6H4SO3H.H2O in benzene under reflux 2 h to give 202 g 1-(3,4-dimethoxyphenyl)-2,3bis (methoxycarbonyl) -4-hydroxy-6,7,8-trimethoxynaphthalene (III). Rats fed with a feed contg. 20 mg% III showed serum cholesterol decrease by

52% and HDL-cholesterol increase by 86%.

104756-71-0P IT

> RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. and cycloaddn. of, with di-Me acetylenedicarboxylate)

RN 104756-71-0 HCAPLUS

CN Benzenemethanol, 6-(dimethoxymethyl)-.alpha.-(3,4-dimethoxyphenyl)-2,3,4trimethoxy- (9CI) (CA INDEX NAME)

- L9 ANSWER 15 OF 21 HCAPLUS COPYRIGHT 1999 ACS
- 1988:221419 HCAPLUS ΜA
- DN 108:221419
- ΤI Hypolipidemic naphthalenedicarboxylate derivatives, processes for their preparation, and their pharmaceutical compositions
- IN Iwasaki, Tameo; Takashima, Kohki
- Tanabe Seiyaku Co., Ltd., Japan PA
- Eur. Pat. Appl., 34 pp. SO

CODEN: EPXXDW

- DTPatent
- English LA

FAN.	CNT 1				
	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	EP 251315	A2	19880107	EP 87-109481	19870701
	EP 251315	A3	19890607		
	EP 251315	B1	19911009		
	R: AT, BE,	CH, DE	, ES, FR, GB,	GR, IT, LI, LU, NL	, SE
	JP 63010746	A2	19880118	JP 86-155416	19860701
	US 4840951	Α	19890620	US 87-64293	19870617
	CA 1294278	A1	19920114	CA 87-540829	19870629
	AT 68172	E	19911015	AT 87-109481	19870701
	ES 2038622	Т3	19930801	ES 87-109481	19870701
PRAI	JP 86-155416	19860	701		
	EP 87-109481	19870	701		
OS	MADDAT 108.2214	19			

MARPAT 108:221419

GI

AB Title compds. I (R1, R2 = OR5, NHR5, NR6R7; one of R1 and R2 may = lower alkoxy; R3, R4 = lower alkoxy; one of R3 and R4 may = H; R5 = substituted alkyl, heterocyclyl, or alkenyl; R6, R7 = H, lower alkyl; ring A may be substituted) are prepd. for use as hypolipidemic agents. Amidation of 1-(3,4-dimethoxyphenyl)-2-methoxycarbonyl-4-benzyloxy-6,7,8-trimethoxy-3-naphthoic acid with isobutylamine using 1-hydroxybenzotriazole and DCC, followed by hydrogenolysis of the benzyl group over Pd/C at 3 kg/cm2 H, gave

(dimethoxyphenyl) (methoxycarbonyl) (isobutylcarbamoyl) hydroxytrimethox ynaphthalene II. At 100 mg/kg orally in rats, II decreased serum cholesterol by 60.0% and increased serum HDL-cholesterol by 99.0%.

IT 104756-71-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and cyclocondensation of, with acetylenedicarboxylate)

RN 104756-71-0 HCAPLUS

CN Benzenemethanol, 6-(dimethoxymethyl)-.alpha.-(3,4-dimethoxyphenyl)-2,3,4-trimethoxy- (9CI) (CA INDEX NAME)

- L9 ANSWER 16 OF 21 HCAPLUS COPYRIGHT 1999 ACS
- AN 1986:572073 HCAPLUS
- DN 105:172073
- TI Naphthalene derivatives and their pharmaceutical compositions
- IN Iwasaki, Tameo; Takashima, Kohki
- PA Tanabe Seiyaku Co., Ltd., Japan
- SO Eur. Pat. Appl., 70 pp.
 - CODEN: EPXXDW
- DT Patent
- LA English

FAN.CNT	1					
PA'	TENT NO.	KIND	DATE	API	PLICATION NO.	DATE
PI EP	188248	A2	19860723	EP	86-100282	19860110
EP	188248	A3	19861217			
EP	188248	B1	19900711			
	R: AT, BE	, CH, DE	, FR, GB, I'	T, LI, I	LU, NL, SE	
IL	77457	A1	19910310	$_{ m IL}$	85-77457	19851226
IL	91117	A1	19910310	IL	85-91117	19851226
NO	8505355	Α	19860711	NO	85-5355	19851230
МО	170760	В	19920824			
МО	170760	С	19921202			
ES	550578	A1	19870516	ES	85-550578	19851230
US	4771072	Α	19880913	US	85-814805	19851230
AU	8551751	A1	19860717	AU	85-51751	19851231
AU	584153	B2	19890518			
JP	61267541	A2	19861127	JP	86-2624	19860108
FI	8600089	Α	19860711	FI	86-89	19860109
	87557	В	19921015			
FI	87557	C	19930125			
HU	42428	A2	19870728	HU	86-90	19860109
HU	196737	В	19890130			
SU	1581217	A3	19900723		86-4013137	19860109
CN	86100090	A	19860820	CN	86-100090	19860110
	1006464	В	19900117			
	261786	A5	19881109		86-286106	19860110
	54441	E	19900715		86-100282	19860110
	557052	A1	19871216	ES	86-557052	19860903
	1577697	A3	19900707		86-4028493	19861113
	4897418	Α	19900130	US	88-144650	19880111
	270529	A5	19890802	DD	88-312249	19880115
	01301652	A2	19891205	JP	88-310355	19881208
	06000724	B4	19940105			
JP	02072136	A2	19900312	JP	88-310353	19881208
JP	02072170	A2	19900312	JP	88-310354	19881208
JP	05049668	B4	19930726			
US	5070103	Α	19911203	US	90-459859	19900102
PRAI JP	85-3090	19850	110			
JP	86-2624	19850	110			
IL	85-77457	19851	226			
US	85-814805	19851	230			
EP	86-100282	19860	110			
US	88-144650	19880	111			
GI						

AB Naphthalene derivs. I (R1 = H, alkoxycarbonyl; R2 = alkoxycarbonyl; R1R2

CH2O2C; R3 or R4 = alkoxy, the other = H, alkoxy; R5-R8 = H, substituent) were prepd. (40 examples) as agents for the treatment or prophylaxis of hyperlipidemia and/or arteriosclerosis. Thus, 2,3,4,5-

Br(MeO)3C6HCH(OMe)2
 in THF was treated with BuLi and 3,4-(MeO)2C6H3CHO to give benzaldehyde
 deriv. II, which reacted with MeO2CC.tplbond.CCO2Me in the presence of
 p-MeC6H4SO3H.H2O to give I (R1 = R2 = CO2Me, R3-R7 = OMe, R8 = H) (III).
 At 20 mg% in the diet of rats, III gave 52% redn. of total serum

IT 104756-71-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and cyclocondensation of, with acetylenedicarboxylate)

cholesterol, and increased serum HDL-cholesterol by 86%.

RN 104756-71-0 HCAPLUS

CN Benzenemethanol, 6-(dimethoxymethyl)-.alpha.-(3,4-dimethoxyphenyl)-2,3,4trimethoxy- (9CI) (CA INDEX NAME)

- L9 ANSWER 17 OF 21 HCAPLUS COPYRIGHT 1999 ACS
- AN 1986:226523 HCAPLUS
- DN 104:226523
- TI Chemical structures of sulfuric acid lignin. IX. Reaction of syringyl alcohol and reactivity of guaiacyl and syringyl nuclei in sulfuric acid solution
- AU Yasuda, Seiichi; Ota, Katsuhito
- CS Fac. Agric., Nagoya Univ., Nagoya, 464, Japan
- SO Mokuzai Gakkaishi (1986), 32(1), 51-8 CODEN: MKZGA7; ISSN: 0021-4795
- DT Journal
- LA English
- AB The behavior of syringyl and guaiacyl nucleus of lignin in H2SO4 was studied by model reaction of syringyl alc. [530-56-3], 3,4,5-trimethoxybenzyl alc. [3840-31-1], vanillyl alc. (I) [498-00-0] and veratryl alc. [93-03-8] with creosol (II) [93-51-6] and II Me ether [494-99-5]; reaction of acetoguaiacone Me ether [91-10-1] with II, condensation of I with various arom. compds.; condensation of apocynol Me ether [5653-65-6] with II and 5-methoxycresol [6638-05-7]; and condensation of propionaldehyde [123-38-6] with II. Based on results from the reaction of I with arom. compds. in 5% H2SO4, the reactivity of

arom. nuclei decreased in the order: syringyl > etherified syringyl > etherified guaiacyl > guaiacyl.

102430-92-2P IT

> RL: FORM (Formation, nonpreparative); PREP (Preparation) (formation of, in model reactions for lignin in sulfuric acid)

102430-92-2 HCAPLUS RN

Phenol, 3-[1-(3,4-dimethoxyphenyl)ethyl]-2,6-dimethoxy-4-methyl- (9CI) CN (CA INDEX NAME)

102415-83-8 IT

RL: RCT (Reactant)

(reaction of, with creosol, in sulfuric acid, as lignin model)

RN102415-83-8 HCAPLUS

Benzene, 1,2,3-trimethoxy-5-methyl-4-[1-(3,4,5-trimethoxyphenyl)ethyl]-CN (9CI) (CA INDEX NAME)

ANSWER 18 OF 21 HCAPLUS COPYRIGHT 1999 ACS L9

AN 1983:612363 HCAPLUS

DN 99:212363

Hydroxy acetals, phthalans, and isobenzofurans therefrom TI

Keay, Brian A.; Plaumann, Heinz P.; Rajapaksa, Dayananda; Rodrigo, ΑU Russell

Guelph-Waterloo Cent. Grad. Work Chem., Univ. Waterloo, Waterloo, ON, N2L 3G1, Can.

so Can. J. Chem. (1983), 61(9), 1987-95 CODEN: CJCHAG; ISSN: 0008-4042

DTJournal

English LA

GI

CS

- AB A general method for the generation of isobenzofuran intermediates is described. Lithiated arom. acetals are converted to hydroxy acetals I (R = substituted Ph, R1-R4 = H, OMe, R2R3 = OCH2O), which are cyclized to isobenzofurans by mild acid treatment through the hydroxyphthalans II (R5 = H, Me). The isobenzofurans generated in situ are trapped by a variety of dienophiles to give the oxabicyclo adducts, e.g., III. The mass spectra and NMR spectra of II and III are discussed.
- RN 87850-24-6 HCAPLUS
 CN Benzenemethanol, 6-(dimethoxymethyl)-2,3,4-trimethoxy-.alpha.-(3,4,5-trimethoxyphenyl)- (9CI) (CA INDEX NAME)

- L9 ANSWER 19 OF 21 HCAPLUS COPYRIGHT 1999 ACS
- AN 1978:169703 HCAPLUS
- DN 88:169703
- TI Reactions of halomagnesium alcoholates of aromatic alcohols with perfluorinated halomagnesium thiophenolates in the presence of ethyl formate
- AU Bogoslovskii, N. V.; Kolbina, N. M.
- CS Perm. Gos. Univ., Perm, USSR
- SO Org. Khim. (1976), 39-43. Editor(s): Lapkin, I. I. Publisher: Permsk. Gos. Univ. im. A. M. Gor'kogo, Perm, USSR. CODEN: 37LPAM
- DT Conference
- LA Russian

AB C6F5MgCl reacted with S to give C6F5SMgCl, which reacted with RCH2OMgBr (R

= Ph, 3,4-Cl2C6H3, .alpha.-naphthyl) and HCO2Et to give 45-55% RCH2SC6F5 (I). I were oxidized with 30% H2O2 to yield 88-98% RCH2SO2C6F5. The analogous reaction of C6F5CHROMgCl [R = Ph, 4-ClC6H4, 4-BrC6H4, 2,4-Cl2C6H3, 3,4-(MeO)2C6H3] (from C6F5MgCl and RCHO) gave 57-81% C6F5CHROH but no sulfides.

IT 66390-45-2P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (prepn. of)

RN 66390-45-2 HCAPLUS

CN Benzenemethanol, .alpha.-(3,4-dimethoxyphenyl)-2,3,4,5,6-pentafluoro-(9CI) (CA INDEX NAME)

L9 ANSWER 20 OF 21 HCAPLUS COPYRIGHT 1999 ACS

AN 1972:126515 HCAPLUS

DN 76:126515

TI Reactions of halometal alcoholates. I. Synthesis of methylhydroxydiarylmethanes

AU Lapkin, I. I.; Belonovich, M. I.; D'yakova, G. F.

CS Perm. Gos. Univ., Perm, USSR

SO Zh. Org. Khim. (1972), 8(2), 292-3 CODEN: ZORKAE

DT Journal

LA Russian

AB RCHMeOMgBr (R = Ph, 2-MeOC6H4, 2- and 4-MeC6H4, 2,5-Me2C6H3, 2,4,6-Me3C6H2) reacted with HCO2Et to form RCHMeBr, which gave the corresponding RCHMeR1 (R1 = hydroxyaryl) in 40-70% yield with 7 R10MgBr.

IT 35770-83-3P 35770-85-5P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of)

RN 35770-83-3 HCAPLUS

CN Phenol, 2-methyl-4-[1-(2,4,6-trimethylphenyl)ethyl]- (9CI) (CA INDEX NAME)

RN 35770-85-5 HCAPLUS

CN Phenol, 5-methyl-2-(1-methylethyl)-4-[1-(2,4,6-trimethylphenyl)ethyl](9CI) (CA INDEX NAME)

L9 ANSWER 21 OF 21 HCAPLUS COPYRIGHT 1999 ACS

AN 1970:89960 HCAPLUS

DN 72:89960

TI Reaction of polyfluoro-substituted aromatic ketones with potassium cyanide

AU Vasilevskaya, T. N.; Badashkeeva, A. G.; Gerasimova, T. N.; Barkhash, V. A.; Vorozhtsov, N. N., Jr.

CS Novosibirsk. Inst. Org. Khim., Novosibirsk, USSR

SO Zh. Org. Khim. (1970), 6(1), 126-32 CODEN: ZORKAE

DT Journal

LA Russian

The vigorous reaction of (C6F5)2CO with KCN in abs. EtOH at 20.degree. gave C6F5H, 2,3,5,6-F4C6HCN (I), C6F5CO2Et (II), 2,3,5,6,4-F4(EtO)C6CO2Et (III), and 2,3,5,6,7-F4(EtO)C6COC6F5 (IV). The compds. were sepd. by gas chromatog. and identified by NMR. The reaction of II with EtONa gave

III.

Refluxing C6F5Br with EtONa in EtOH gave 2,3,5,6,4-F4(EtO)C6Br (V) which was converted to its Grignard compd. and reacted with C6F5CHO to give 2,3,5,-6,4-F4(EtO)C6CH(OH)C6F5, which on oxidn. with CrO3 gave IV. The reaction of C6F5COPh with KCN in EtOH at 75.degree. gave C6F5H, I, and 2,3,5,6,4-F4(EtO)C6COPh (VI). Reacting V with Mg and PhCHO in abs. Et2O gave 2,3,5,6,4-F4(EtO)-C6CH(OH)Ph which was oxidized to VI. The reaction of C6F5-COMe with KCN in EtOH at 60-70.degree. gave C6F5H, I, AcOEt, 2,3,5,6-F4C6HC(:NH)OEt (VII), 3,5,6,2-F3(EtO)C6HCN, and 2,3,5,6,4-F4(EtO)C6COMe (VIII). Treating V with Mg and Ac2O gave VIII. The treatment of VII with HCl in Et2O gave 2,3,5,6-F4C6HCONH2.

IT 28293-48-3P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (prepn. of)

RN 28293-48-3 HCAPLUS

CN Benzhydrol, 4-ethoxy-2,2',3,3',4',5,5',6,6'-nonafluoro- (8CI) (CA INDEX NAME)

Benzoquinone structures

L4 ANSWER 1 OF 4 HCAPLUS COPYRIGHT 1999 ACS

AN 1999:9803 HCAPLUS

TI Preparation of phenoxyakanoates as thyroid hormone receptor .beta. agonists

IN Scanlan, Thomas S.; Chellini, Grazia; Yoshihara, Hikari; Apriletti,
James;

Baxter, John D.; Ribeiro, Ralff C. J.

PA The Regents of the University of California, USA

SO PCT Int. Appl., 45 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE

PI WO 9857919 A1 19981223 WO 98-US11758 19980608

CO2H

W: AU, CA, JP, KP, KR

RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT SE

PRAI US 97-877792 19970618 GI

Me Me | I

AB R3OZ1CR1R2Z2O(CH2)nCO2R [I; R = H or (cyclo)alkyl; R1,R2 = H or alkyl; 1 of R1,R2 = H and the other = OH; R1R2 = O; R3 = H, (cyclo)alkyl, acyl; Z1 = (un)substituted 1,4-phenylene; Z2 = (un)substituted 3,5-dimethyl-4,1-phenylene] were prepd. Thus, 4-bromo-2-isopropylanisole was condensed with 2,6-dimethyl-4-methoxybenzaldehyde (prepn. each given) and the product converted in 4 steps to title compd. II. Data for biol. activity of I were given.

IT 218431-20-0P 218431-21-1P 218431-24-4P 218431-25-5P 218431-26-6P

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of phenoxyakanoates as thyroid hormone receptor .beta. agonists)

RN 218431-20-0 HCAPLUS

CN INDEX NAME NOT YET ASSIGNED

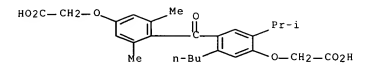
RN 218431-21-1 HCAPLUS CN INDEX NAME NOT YET ASSIGNED

RN 218431-24-4 HCAPLUS CN INDEX NAME NOT YET ASSIGNED

RN 218431-25-5 HCAPLUS CN INDEX NAME NOT YET ASSIGNED

RN 218431-26-6 HCAPLUS CN INDEX NAME NOT YET ASSIGNED





IT 214544-31-7P 218431-17-5P 218431-19-7P

218431-22-2P 218431-23-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
 (prepn. of phenoxyakanoates as thyroid hormone receptor
 .beta. agonists)

RN 214544-31-7 HCAPLUS

CN Methanone, (4-methoxy-2,6-dimethylphenyl) [4-methoxy-3-(1-methylethyl)phenyl] - (9CI) (CA INDEX NAME)

RN 218431-17-5 HCAPLUS

CN INDEX NAME NOT YET ASSIGNED

RN 218431-19-7 HCAPLUS

CN INDEX NAME NOT YET ASSIGNED

RN 218431-22-2 HCAPLUS

CN INDEX NAME NOT YET ASSIGNED

$$\begin{array}{c|c} \text{MeO} & \text{Me} & \text{O} \\ \hline \\ \text{Me} & \text{n-Bu} \\ \end{array} \\ \begin{array}{c} \text{Pr-i} \\ \text{OMe} \\ \end{array}$$

RN 218431-23-3 HCAPLUS

CN INDEX NAME NOT YET ASSIGNED

L4 ANSWER 2 OF 4 HCAPLUS COPYRIGHT 1999 ACS

AN 1998:617873 HCAPLUS

DN 129:302827

TI An efficient substitution reaction for the preparation of thyroid hormone analoges

AU Yoshihara, Hikari A. I.; Chiellini, Grazia; Mitchison, Timothy J.; Scanlan, Thomas S.

CS Department of Cellular and Molecular Pharmacology, University of California, San Francisco, CA, 94143-0450, USA

SO Bioorg. Med. Chem. (1998), 6(8), 1179-1183 CODEN: BMECEP; ISSN: 0968-0896

PB Elsevier Science Ltd.

DT Journal

LA English

AB The substitution of the sterically hindered carbon of the potent thyroid hormone agonist, GC-1, was effected by a reaction based on the solvolysis of the benzylic hydroxyl group. The reaction was found to proceed in high yield with a variety of nucleophiles including alcs., thiols, allyl silanes and electron-rich arom. compds., providing a convenient route to the synthesis of new thyroid hormone analogs.

IT 214544-31-7P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
 (prepn. of thyroid hormone analoges via substitution
 reaction)

RN 214544-31-7 HCAPLUS

CN Methanone, (4-methoxy-2,6-dimethylphenyl) [4-methoxy-3-(1methylethyl)phenyl] - (9CI) (CA INDEX NAME)

IT 214544-32-8P 214544-34-0P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (prepn. of thyroid hormone analoges via substitution
 reaction)

RN 214544-32-8 HCAPLUS

CN Methanone, [2-butyl-4-methoxy-3-(1-methylethyl)phenyl] (4-methoxy-2,6dimethylphenyl) - (9CI) (CA INDEX NAME)

RN 214544-34-0 HCAPLUS

CN Methanone, (4-methoxy-2,6-dimethylphenyl)[4-methoxy-3-(1-methylethyl)-2-

(1 methylpropyl)phenyl] - (9CI) (CA INDEX NAME)

L4 ANSWER 3 OF 4 HCAPLUS COPYRIGHT 1999 ACS

AN 1984:584212 HCAPLUS

DN 101:184212

TI Comparative effects of thyroid hormone analogs on the activities of brain and liver mitochondria and nuclei in thyroidectomized rats

AU Dembri, A.; Michel, R.; Michel, O.; Belkhiria, M.; Jorgensen, E. C.

CS Coll. France, Paris, 75231, Fr.

SO Mol. Cell. Endocrinol. (1984), 37(2), 223-32 CODEN: MCEND6; ISSN: 0303-7207

DT Journal

LA English

AB Several thyroid hormone analogs were tested for thyromimetic

activity on rat brain and liver subcellular organelles. The compds. were administered immediately after thyroidectomy to 90 g male rats for 10 days, by daily s.c. injection. In cerebral cortex and liver, the activities of mitochondrial succinate cycochrome c reductase [9028-10-8] and .alpha.-glycerophosphate dehydrogenase [9075-65-4] and nuclear RNA polymerase [9014-24-8] were measured. Brain mitochondrial enzymes were unchanged in thyroidectomized (Tx) and in Tx-treated rats, whereas the activities of these enzymes in liver mitochondria were partially restored by the treatments. RNA polymerase I activity in brain and liver dropped significantly 10 days after thyroidectomy and daily injection of thyroid hormones or analogs maintained the nuclear activity at a normal level. Correlation between the structure of thyroid hormone analogs and their subcellular effects is in good agreement with previous binding and in vivo studies. Enzyme activities stimulated by T3 [6893-02-3] were lowered by replacing the T3 side-chain by an acetic acid group or by substituting the bridged O atom by atom by CO. In contrast, the activity was enhanced by substituting I with a 3' iso-Pr group. Although less active than I, the 3,5-di-Me substituents may be introduced without a complete loss of nuclear activity.

IT 92814-41-0

RL: BAC (Biological activity or effector, except adverse); BIOL (Biological study)

(thyromimetic activity of, structure in relation to)

RN 92814-41-0 HCAPLUS

CN Benzeneacetic acid, 4-[4-hydroxy-3-(1-methylethyl)benzoyl]-3,5-diiodo-(9CI) (CA INDEX NAME)

L4 ANSWER 4 OF 4 HCAPLUS COPYRIGHT 1999 ACS

AN 1982:518486 HCAPLUS

DN 97:118486

TI Methyl 3,5-diiodo-4-(3-isopropyl-4-methoxybenzoyl)benzoate

AU Cody, Vivian; Cheung, Ellen; Jorgensen, Eugene C.

CS Med. Found. Buffalo, Inc., Buffalo, NY, 14203, USA

SO Acta Crystallogr., Sect. B (1982), B38(8), 2270-2 CODEN: ACBCAR; ISSN: 0567-7408

DT Journal

LA English

AB The title compd. is orthorhombic, space group Iba2, with a 20.998(3), b 24.002(4), and c 8.032(1) .ANG.; Z = 8 for dc = 1.85; R = 6.6%. The conformation of the di-Ph ketone bridge is skewed and the iso-Pr group distally oriented, as is obsd. for many thyroid hormone analog structures. There is a short I...O intermol. contact between I(5) and

carbonyl O [3.17(10) .ANG.]. At. coordinates are given.

IT 82897-04-9

the

RL: PRP (Properties)

(structure of)

CN

RN 82897-04-9 HCAPLUS

Benzoic acid, 3,5-diiodo-4-[4-methoxy-3-(1-methylethyl)benzoyl]-, methyl
ester (9CI) (CA INDEX NAME)

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